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* * * * * * * * * * * Welcome to STN International
      * * * * * * * * * * * * * STN Columbus * * * * * * * * * * *
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                                                        13
                                                                          chain nodes :
11 12 13 14 15
ring nodes :
1 2 3 4 5 6 7 8 9 10
chain bonds :
1-15 4-13 8-11 11-12 13-14
ring bonds :
1-2 1-6 2-3 3-4 4-5 4-7 5-6 5-10 7-8 8-9 9-10
exact/norm bonds :
1-2 1-6 1-15 2-3 3-4 4-5 4-7 4-13 5-6 5-10 7-8 8-9 8-11 9-10 11-12
13-14
isolated ring systems :
containing 1 :
Match level :
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom
11:CLASS 12:Atom 13:CLASS 14:Atom 15:CLASS
=> s 11 sam
             8 SEA SSS SAM L1
L2
=> s 11 full
L3
            96 SEA SSS FUL L1
=> file caplus
=> s 13
           10 L3
L4
=> s 14 and pd< jan 2003
      23690593 PD< JAN 2003
                (PD<20030100)
L5
            5 L4 AND PD< JAN 2003
=> dis 15 1-5 bib abs hitstr
    ANSWER 1 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
L5
AN
    2003:613051 CAPLUS Full-text
DN
    140:339511
    Synthetic study of nakadomarin A
AU
    Nishida, Atsushi; Nagata, Toshiaki; Kano, Takuya; Ono, Kouji; Nakagawa,
     Masako
    Graduate School of Pharmaceutical Sciences, Chiba University, Japan
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SO Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (2001), 43rd, 617-622

CODEN: TYKYDS

- PB Nippon Kagakkai DT Journal
- DT Journal
- LA Japanese
- AB A symposium report. The alkaloid, nakadomarin A has been isolated from the Okinawan marine sponga Amphimedon sp. by Kobayashi in 1997. Although nakadomarin A is a member of manzamine family, it has a unique structure consisting of unprecedented 8/5/5/5/15/6 ring system containing a furan ring. Because of our interest to its unique ring system and also a need for enough quantity for full biol. test, we started the synthesis of nakadomarin A our two synthetic approaches to the central core ring system of nakadomarin A were reported. Route A consists of Diels-Alder reaction of chiral dienophile and siloxydiene to give a 6/6 AB ring system, followed by conversion to 5/5/6 ABD ring system by ozonolysis and intramol. aldol reaction. Route B is an efficient route for the synthesis of racemic ABCD ring system. A key step was intramol. cyclization of a furan ring to acyl iminium cation. Ring closing metathesis of a diene to construct ring F was also discussed.
- IT 221526-51-8P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
 (Reactant or reagent)
 - (synthetic study of nakadomarin A)
- RN 221526-51-8 CAPLUS
- CN 1,6(2H,5H)-Tsoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aS,8aS)-(CA INDEX NAME)

Absolute stereochemistry.

- L5 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2003:280654 CAPLUS Full-text
- DN 139:245967
- TI Practical synthesis of a 3, 4, 4a, 5, 8, 8a-hexahydro-2H-isoquinoline-1,6-dione ring system by the Diels-Alder reaction of an optically active dienophile, a 5,

6-dihydro-1H-pyridin-2-one derivative, with siloxydiene

- AU Nakagawa, Masako; Uchida, Hideharu; Ono, Koji, Kimura, Yoshiyuki; Yamabe, Mariko; Watanabe, Takeshi; Tsuji, Riichiro; Akiba, Masakatsu; Terada, Yukiyoshi; Nagaki, Dai; Ban, Sachiko; Miyashita, Naoki; Kano, Takuya; Theeraladanon, Chumpol; Hatakeyama, Keisuke; Arisawa, Mitsuhiro; Nishida, Atsushi
- CS Graduate School of Pharmaceutical Sciences, Chiba University, Chiba, 263-8522, Japan
- SO Heterocycles (2003), 59(2), 721-733 CODEN: HTCYAM; ISSN: 0385-5414
- PB Japan Institute of Heterocyclic Chemistry

- DT Journal
- LA English
- OS CASREACT 139:245967
- AB An efficient method for preparing chiral 3-substituted-5, 6-dihydro-IHpyridin-2-one (I) in large scale, based on a modification of our previous method, is described. The large scale Diels-Alder reaction of I with siloxydiene to synthesize hexahydroisoquinoline-1,6-dione, which is a key intermediate for the synthesis of manzamine alkaloids, was also studied.
- T 221526-51-8P 221526-52-9P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
 - (Reactant or reagent)

 (multi-step preparation of chiral substituted dihydropyridinone and its
 Diels-Alder reaction with siloxydiene to hexahydroisoguinolinedione as
- Diels-Alder reaction with siloxydiene to hexahydroisoquinolinedione as key intermediate for synthesis of manzamine alkaloids)
- RN 221526-51-8 CAPLUS
- CN 1,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aS,8aS)-(CA INDEX NAME)

Absolute stereochemistry.

- RN 221526-52-9 CAPLUS
- CN 1,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aR,8aR)-(CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L5 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 1999:776314 CAPLUS Full-text
- DN 132:279387
- TI Total synthesis of manzamine A and related compounds
- AU Uchida, Hideharu; Kimura, Yoshiyuki; Yamabe, Mariko; Nishida, Atsushi; Nakagawa, Masako

- CS Faculty of Pharmaceutical Sciences, Chiba University, Japan SO Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (1999), 41st, 67-72 CODEN: TYKYDS
- PB Nippon Kagakkai
- DT Journal
- LA Japanese

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AR Manzamine alkaloids constitute a novel family of several marine sponge metabolites that exhibit significant cytotoxic activity against leukemia and antibiotic activity. The unprecedented structures of these highly functionalized heterocyclic ring system and remarkable biol. properties have attracted much attention as a challenging synthetic targets. While the simplest congener manzamine C and related compds. have been previously synthesized by us and Langlois' group, the more complex manzamine A has been more challenging target. Quite recently, Winkler, Martin and their coworkers have succeeded in total synthesis of manzamine A and its related compds. The authors have also been interested in developing efficient routes to tetraazacyclic intermediate (I) based on the initial construction of tricyclic intermediate (II) by an intermol. Diels-Alder reaction of functionalized dihydropyridinone (III) as a dienophile with siloxydiene, leading to the construction of a cis relationship in the central AB ring system of this unique structure. Using this strategy, the authors could synthesize advanced key intermediates (IV) and (V). Details of the synthesis and further conversion to ircinal A and manzamine A will be discussed.

IT 221526-51-8P 221526-52-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(total synthesis of manzamine A and related compds.)

- RN 221526-51-8 CAPLUS
- CN 1,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aS,8aS)-(CA INDEX NAME)

Absolute stereochemistry.

- RN 221526-52-9 CAPLUS
- CN 1,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aR,8aR)-(CA INDEX NAME)

L5 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN

AN 1999:313230 CAPLUS Full-text

DN 131:228862

TI Synthetic studies on optically active ircinal A

AU Uchida, Hideharu; Takezawa, Emiko; Kawate, Tomohiko; Nishida, Atsushi; Nakagawa, Masako

CS Faculty of Pharmaceutical Sciences, Chiba University, Japan

SO Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (1998), 40th, 601-606

CODEN: TYKYDS PB Nippon Kagakkai

DT Journal

LA Japanese

GT Japanese

II

- AB Ircinal A was isolated from Okinawan marine sponges by Kobayashi. Ircinal A and manzamine A have been the subject of recent synthetic investigations owing its unique mol. structure and remarkable biol. properties including antitumor and antibacterial activities. The authors have already accomplished a synthesis of key tetracyclic intermediate (ABCD ring) for ircinal A and manzamine A in a racemic form. For the synthesis of optically active ircinal A and manzamine A, several optically active dienophiles were prepared and subjected to Diels-Alder reaction with Danishefsky diene. The D-A adduct, obtained by the reaction of dienophile I, was successfully converted to the tetracyclic intermediate II, in an optically active form.
- IT 221526-51-8P 221526-52-9P 244074-45-1P 241074-46-2P 244074-51-9P 244074-52-0P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (synthetic studies on optically active ircinal A and manzamine A)
- RN 221526-51-8 CAPLUS
- CN 1,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-

oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aS,8aS)-(CA INDEX NAME)

Absolute stereochemistry.

- RN 221526-52-9 CAPLUS
- CN 1,6(2R,58)-Isoquinolinedione, 8a-[((4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl|-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aR,8aR)-(CA INDEX NAME)

Absolute stereochemistry.

- RN 244074-45-1 CAPLUS
- CN 3-Oxazolidinecarboxylic acid, 4-[[(4aS,8aS)-2,3,4,4a,5,6-hexahydro-1,6-dioxo-2-(phenylsulfonyl)-8a(1H)-isoquinolinyl]methyl]-2,2-dimethyl-,1,1-dimethylethyl ester, (4R)- (CA INDEX NAME)

- RN 244074-46-2 CAPLUS
- CN 1,6(2H,5H)-Isoquinolinedione, 2-[(4-chlorophenyl)sulfonyl]-8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-,(4aS,8aS)- (CA INDEX NAME)

Absolute stereochemistry.

RN 244074-51-9 CAPLUS

CN 3-Oxazolidinecarboxylic acid, 4-[[(4aR,8aR)-2,3,4,4a,5,6-hexahydro-1,6-dioxo-2-(phenylsulfonyl)-8a(1H)-isoquinolinyl]methyl]-2,2-dimethyl-,1,1-dimethylethyl ester, (4R)- (CA INDEX NAME)

Absolute stereochemistry.

- RN 244074-52-0 CAPLUS
- CN 1,6(2H,5H)-Tsoquinolinedione, 2-[(4-chlorophenyl)sulfonyl]-8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-,(4aR,8aR)- (CA INDEX NAME)

- L5 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 1999:83174 CAPLUS Full-text
- DN 130:252516
- $\ensuremath{\text{TI}}$. An efficient access to the optically active manzamine tetracyclic ring system
- AU Uchida, Hideharu; Nishida, Atsushi; Nakagawa, Masako
- CS Faculty of Pharmaceutical Sciences, Chiba University, Chiba, 263-8522,

Japan

SO Tetrahedron Letters (1999), 40(1), 113-116 CODEN: TELEAY: ISSN: 0040-4039

PB Elsevier Science Ltd.

DT Journal

LA English

OS CASREACT 130:252516

- AB The highly stereoselective synthesis of the optically active tetracyclic core I of Manzamine A was achieved via the Diels-Alder reaction of dihydropyridinone II, derived from L-serine, with siloxydienes, followed by sequential new and conventional pathways.
- IT 221526-51-8P 221526-52-9P

Ι

- RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 - (efficient access to optically active manzamine tetracyclic ring system)
- RN 221526-51-8 CAPLUS
- I,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aS,8aS)-(CA INDEX NAME)

Absolute stereochemistry.

RN 221526-52-9 CAPLUS

CN 1,6(2H,5H)-Tsoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aR,8aR)-(CA INDEX NAME)

RE.CNT 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 14 not 15 L6 5 L4 NOT L5

=> dis 16 1-5 bib abs fhitstr

- L6 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2008:232071 CAPLUS Full-text
- DN 148:440269
- TI 1H-Pyrazolo[3,4-g]hexahydro-isoquinolines as selective glucocorticoid receptor antagonists with high functional activity
- AU Clark, Robin D.; Ray, Nicholas C.; Williams, Karen; Blaney, Paul; Ward, Stuart; Crackett, Peter H.; Hurley, Christopher; Dyke, Hazel J.; Clark, David E.; Lockey, Peter; Devos, Rene; Wong, Melanie; Porres, Soraya S.; Bright, Colin P.; Jenkins, Robert E.; Belanoff, Joseph
- CS Corcept Therapeutics, Menlo Park, CA, 94025, USA
- SO Bioorganic & Medicinal Chemistry Letters (2008), 18(4), 1312-1317 CODEN: BMCLE8; ISSN: 0960-894X
- PB Elsevier Ltd.
- DT Journal
- LA English
- OS CASREACT 148:440269
- AB Addition of the 4-fluorophenylpyrazole group to the previously described 2azadecalin glucocorticoid receptor (GR) antagonist I resulted in significantly enhanced functional activity. SAR of the bridgehead substituent indicated that whereas groups as small as Me afforded high GR binding, GR functional activity was enhanced by larger groups such as benzyl, substituted ethers, and aminoalkyl derivs. GR antagonists with binding and functional activity comparable to mifepristone were discovered (e.g., 52: GR binding Kl 0.7 nM, GR reporter gene functional Ki 0.6 nM) and found to be highly selective over other steroid receptors. Analogs 43 and 45 had >50% oral bioavailability in the dog.
- IT 864973-54-6

RL: RCT (Reactant); RACT (Reactant or reagent)

(1H-pyrazolo[3,4-g]hexahydro-isoquinolines as selective glucocorticoid receptor antagonists)

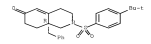
- RN 864973-54-6 CAPLUS
- CN 6(2H)-Isoquinolinone, 2-[[4-(1,1-dimethylethyl)phenyl]sulfonyl]-1,3,4,7,8,8a-hexahydro-8a-(phenylmethyl)-, (8aR)- (CA INDEX NAME)

$$\bigcap_{Ph} \bigcap_{Ph} \bigcap_{Ph}$$

RE.CNT 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2007:1051323 CAPLUS Full-text
- DN 147:534024
- TI 2-Benzenesulfonyl-8a-benzyl-hexahydro-2H-isoquinolin-6-ones as selective glucocorticoid receptor antagonists
- AU Clark, Robin D.; Ray, Nicholas C.; Blaney, Paul; Crackett, Peter H.; Hurley, Christopher; Williams, Karen; Dyke, Hazel J.; Clark, David E.; Lockey, Peter M.; Devos, Rene; Wong, Melanie; White, Anne; Belanoff, Joseph K.
- CS Corcept Therapeutics, Menlo Park, CA, 94025, USA
- SO Bioorganic & Medicinal Chemistry Letters (2007), 17(20), 5704-5708 CODEN: BMCLE8; ISSN: 0960-894X
- PB Elsevier Ltd.
- DT Journal
- LA English
- OS CASREACT 147:534024
- AB The 2-azadecalin ring system was evaluated as a scaffold for the preparation of glucocorticoid receptor (GR) antagonists. High affinity, selective GR antagonists were discovered based on a hypothetical binding mode related to the steroidal GR antagonist RU-43044. 2-Benzenesulfonyl substituted 8a-benzyl hexahydro-2H-isoquinolin-6-ones exemplified by (R)-37 had low nanomolar affinity for GR with moderate functional activity (200 nM) in a reporter gene assay. These compds. were devoid of affinity for other steroidal receptors (ER, AR, MR, and FR). Analogs based on an alternative putative binding mode (CP-like) were found to be inactive.
- IT 864973-54-6P
 - RL: PAC (Pharmacological activity); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (2-benzenesulfonyl-8a-benzyl-hexahydro-2H-isoquinolin-6-ones as selective glucocorticoid receptor antagonists)
- RN 864973-54-6 CAPLUS
- CN 6(2H)-Isoquinolinone, 2-[[4-(1,1-dimethylethyl)phenyl]sulfonyl]-1,3,4,7,8,8a-hexahydro-8a-(phenylmethyl)-, (8aR)- (CA INDEX NAME)

Absolute stereochemistry.



RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD

- L6 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2005:1021750 CAPLUS Full-text
- DN 143:306309
- TI Preparation of triazacyclopenta[b]naphthalene derivatives as modulators of glucocorticoid receptor
- IN Clark, Robin D.; Ray, Nicholas C.; Blaney, Paul M.; Hurley, Christopher A.; Williams, Karen
- Corcept Therapeutics, Inc., USA PA
- SO PCT Int. Appl., 160 pp. CODEN: PIXXD2
- DT Patent
- LA English

FAN.CNT 1																				
						KIND DATE				APPLICATION NO.										
PI		2005																		
PI	WU								WO 2005-US8049 BA, BB, BG, BR, BW,											
		W :							DK,											
									IL,											
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		DW.							MZ,										ΔW	
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	ΔII	2005		A1 20050922					ΔII 2	005-	2224	21		20050309						
			A1 20050922																	
		1735		A1 20061227																
		1735308			B1 20080910															
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			IS,	IT,	LI,	LT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR				
	CN	1010	A	A 20070829				CN 2	005-	8001	1481		20050309							
			T 20071011										20050309							
	AT	4079	T	T 20080915				AT 2	005-	7252	95		20050309							
	ES	2313	Т3	T3 20090301				ES 2	005-	7252	95		20050309							
	KR	2007	0296	84		A	A 20070314				KR 2	006-	7209	88		20061009				
				A 20070615			IN 2006-CN3745													
	US	2007	A1	A1 20071206				US 2007-591884						20070507						
PRAI		2004																		
		2005																		
os	CASREACT 143:306309; MARPAT 143:306309																			
GI																				

AB Title compds. I [Il and L2 independently = a bond, O, S, etc.; A =
 (un)substituted 5-6 membered heterocycloalkyl or heteroaryl; R1 = H,
 (un)substituted alkyl, heteroalkyl, etc.; R2 = (un)substituted alkyl,
 heteroalkyl, cycloalkyl, etc.] and their pharmaceutically acceptable salts,
 are prepared and disclosed as modulators of glucocorticoid receptor. Thus, II
 was prepared by cyclization of (S)-Ba-benzyl-2-(4-tert-butyl-benzenssulfonyl 7-[1-hydroxy-meth-(2)-ylidene]-I,3,4,7,8,8a-hexahydro-2H-isoquinolin-6-one
 (preparation given) with hydrazine hydrate. The activity of I was evaluated
 in glucocorticoid receptor binding assay and it was revealed that selected
 compds. of the invention displayed Ic50 values in the range of 10 up to 100 nm
 and others below 10 nM. Pharmaceutical compns. comprising I are disclosed.

II 861629-5a-1

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of triazacyclopenta[b]naphthalene derivs. as modulators of
qlucocorticoid receptor)

RN 861629-54-1 CAPLUS

CN 6(2H)-Isoquinolinone, 2-[[4-(1,1-dimethylethyl)phenyl]sulfonyl]-1,3,4,7,8,8a-hexahydro-8a-(phenylmethyl)- (CA INDEX NAME)

$$0 = \lim_{n \to \infty} \lim_{n \to \infty} \frac{1}{n} = \lim_{n \to \infty}$$

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2005:696879 CAPLUS Full-text
- DN 143:193917
- TI Preparation of azadecalin derivatives as glucocorticoid receptor modulators
- IN Clark, Robin D.; Ray, Nicholas C.; Blaney, Paul; Hurley, Christopher; Williams, Karen; Hunt, Hazel; Clark, David
- PA Corcept Therapeutics, Inc., USA
- SO PCT Int. Appl., 105 pp.
- CODEN: PIXXD2
- DT Patent

LA English

FAN.			KIND		DATE		APPLICATION NO.												
PI	WO 20											20050110							
	W	E AE, CN, GE, LK, NO, TJ, W: BW, AZ, EE,	AG, CO, GH, LR, NZ, TM, GH, BY, ES,	AL, CR, GM, LS, OM, TN, GM, KG,	AM, CU, HR, LT, PG, TR, KE, KZ,	AT, CZ, HU, LU, PH, TT, LS, MD, GB,	AU, DE, ID, LV, PL, TZ, MW, RU, GR,	AZ, DK, IL, MA, PT, UA, MZ, TJ, HU,	BA, DM, IN, MD, RO, UG, NA, TM, IE,	DZ, IS, MG, RU, US, SD, AT, IS,	EC, JP, MK, SC, UZ, SL, BE, IT,	EE, KE, MN, SD, VC, SZ, BG, LT,	EG, KG, MW, SE, VN, TZ, CH, LU,	ES, KP, MX, SG, YU, UG, CY, MC,	FI, KR, MZ, SK, ZA, ZM, CZ, NL,	GB, KZ, NA, SL, ZM, ZW, DE, PL,	GD, LC, NI, SY, ZW, AM, DK, PT,	SM	
	CA 25	MR, 052064 52419	SN,	TD, A1 A1	TD, TG A1 20050804 A1 20050804				AU 2	005- 005-	2064 2552	97 419		20050110					
	EP 17	1761497			В1	20080903			EP 2005-711316 DK, EE, ES, FI, FR,										
	K	IS,	IT,		LT,			NL,											
	JP 20 AT 40 ES 23	7122		T		2008		JP 2006-549454 AT 2005-711316 ES 2005-711316						20050110					
	NO 20 CN 10	060034 111997	56 0		A A		2006 2008	0926 0206	NO 2006-3456 CN 2005-80004074						20060726 20060804				
			20070118 1 20070830 1 20090116				US 2007-596998					20070308							
	RAI US 2004-535460P WO 2005-US607 S CASREACT 143:193917					W 20050110													

OS CASREACT 143:193917; MARPAT 143:19391'

AB Title compds. I [L2-4 = bond, alkylene, etc.; R1 = absent, H, alkyl, heteroalkyl, etc.; R2 = :0, :N-alkoxy, divalent alkylidene, etc.; R3-4 = alkyl, heteroalkyl, cycloalkyl, etc.;] are prepared For instance, II is prepared in several steps from 1-benzyl-4-oxopiperidine-3-carboxylic acid Me

ester•HCl, benzyl bromide, Me vinyl ketone and 4-methoxybenzyl bromide. I are glucocorticoid receptor modulators [no data].

IT 956913-48-7

RL: PRPH (Prophetic)

(Preparation of azadecalin derivatives as glucocorticoid receptor modulators)

RN 956913-48-7 CAPLUS

CN 6(2H)-Isoquinolinone, 2-[[4-(1,1-dimethylethyl)phenyl]sulfonyl]1,3,4,7,8,8a-hexahydro-8a-[[4-(4-morpholinyl)phenyl]methyl]- (CA INDEX NAME)

- L6 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2004:335759 CAPLUS Full-text
- DN 141:38762
- TI Asymmetric total synthesis of (-)-nakadomarin A
- AU Ono, Koji; Nakagawa, Masako; Nishida, Atsushi
- CS Graduate School of Pharmaceutical Sciences, Chiba University, Inage-ku, Chiba-shi, 263-8522, Japan
- SO Angewandte Chemie, International Edition (2004), 43(15), 2020-2023 CODEN: ACIEF5; ISSN: 1433-7851
- PB Wiley-VCH Verlag GmbH & Co. KGaA
- DT Journal
- LA English
- OS CASREACT 141:38762
- GI

AB A key intermediate in the first asym. synthesis of the marine alkaloid (-)nakadomarin A (I), isolated from the marine sponge Amphimedon sp., was the
optically active hydroisoquinoline II. Two sep. ring-closing-metathesis
reactions were crucial to the construction of the 15- and 8-membered rings.
II 201526-51-89

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(asym. total synthesis of (-)-nakadomarin A)

RN 221526-51-8 CAPLUS

CN 1,6(2H,5H)-Isoquinolinedione, 8a-[[(4R)-2,2-dimethyl-3-(phenylsulfonyl)-4-oxazolidinyl]methyl]-3,4,4a,8a-tetrahydro-2-(phenylsulfonyl)-, (4aS,8aS)-(CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

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